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Martin Barl 9/10/97

Martin Barl 9/10/97

PI = Signature Date

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INTRODUCTION

<u>Subject</u>: The research supported by this award is designed to address the need for the discovery and development of new classes of antifungal drugs. As eucaryotic organisms, fungal pathogens provide limited targets for inhibitory or lethal compounds that do not also have deleterious effects on the host. One target that has been exploited successfully in the development of antifungal compounds is the sterol biosynthetic pathway. The drugs that are currently available for intervention in fungal sterol biosynthesis or function are becoming less effective thus making the search for new compounds and sites of action mandatory. Our work proposes to explore previously uncharacterized steps in sterol biosynthesis as possible new sites for new drug discovery. The specific steps in our approach will utilize primary discovery in *Saccharomyces cerevisiae* to address the largely unexplored pathway in *Candida albicans*, a common fungal pathogen.

<u>Purpose</u>: The major purpose of this work is to address the critical rise in fungal infections noted over the last decade (1). A significant amount of the increase has been due to medical advances in other areas that permit entry of opportunistic fungal pathogens into the human body. Included are new surgical procedures and technologies for artificial structural replacements, increased incidence of organ transplants made possible by the development of methods to disarm the immune system, and chemotherapies which are immunosuppressive. In addition, the rise in disease states, especially the situation posed by HIV, has added to the overall problem.

The array of antifungal compounds currently available to deal with this problem is limited and of decreasing efficacy. This has spurred intensive research in the development of new compounds. The primary classes of drugs in use include the polyenes and the azoles. The former have been in use for many years and function by direct interaction with membrane sterol. The result of this interaction is the formation of pores through which leakage of intracellular components occurs leading to cell death (2). Specificity for fungal cells is conferred by the enhanced binding efficiency to ergosterol, the fungal membrane sterol, compared to cholesterol. The binding specificity, however, is not absolute and interaction with host membrane sterol does lead to serious side effects. For example, amphotericin B, a systemically employed polyene, is toxic to renal tissue. Coupled with the emergence of resistance (3, 4), these side effects limit the use of this group of drugs.

The azole antifungals are a newer class of antifungals that are available in many species and in formulations for topical and systemic applications. These drugs are fungistatic thus making their effectiveness in situations where the host is immuno-compromised problematic over the long term. Azoles function by inhibiting the cytochrome P450-mediated removal of the C-14 methyl group from the ergosterol precursor, lanosterol. The azoles are significantly more active against the fungal enzyme as compared to the human version of the C-14 demethylase.

Widespread use of azoles for a number of fungal infections has led to significant and increasing incidence of resistance (5, 6). Within the last two years alone there have been many reports in the literature documenting azole resistance in *C. albicans* (7-11) as well as in other species of *Candida*(12, 13). Furthermore, resistance has also appeared in several other pathogenic fungi including species of *Crytococcus* (14, 15), *Histoplasma* (16), and *Aspergillus* (17). Recent work in *C. albicans* has defined the mechanisms by which resistance occurs. Mutations in and increases in expression of the gene (*ERG11*) for the C-14 demethylase enzyme, the azole target, have been implicated in some resistant isolates (8, 11). Also, over expression of the small molecule efflux systems such as the ABC (ATP-binding cassette) transporter family and major facilitator family of pumps (7, 9, 10) have been identified as resistance mechanisms. Thus, it appears that the efficacy of the azoles is diminishing and that the arsenal of available antifungals will be further reduced in many infections.

Of particular interest to the US Army Defense Women's Health Research Program, are fungal infections unique to women. The most common such afflictions are vaginal infections caused primarily by *C. albicans*. Vaginal yeast infections will occur in three quarters of the female population at least once during their lifetime (18). Although not considered life-threatening, this infection results in sufficient discomfort that time from work and lost efficiency frequently result. This condition is exacerbated under conditions where medical attention is not readily available and extended stays in the field are required. The most often prescribed treatments for yeast vaginitis are polyene and azole drugs. Also of interest to the Army, although not specific to women, are deep tissue infections caused by accidents or wounds and burns, both of which are commonly colonized by opportunistic fungal pathogens.

Scope: The basic approach of this research is to identify steps in sterol biosynthesis that might be new targets for the development of novel antifungal compounds. This work closely follows our work in *S. cerevisiae*, which though not a pathogen, is an excellent model system for eucaryotic cell studies, in general, and sterol biosynthesis studies, in particular. To that end we have isolated and characterized the genes of many of the steps in the yeast sterol pathway. One of those steps, the transmethylation at position C-24 (*ERG6*), is of particular interest to us and further exploration of this gene in *C. albicans* is one of the aims of this project.

Mutations in the *ERG6* gene of *S. cerevisiae* were originally isolated based on resistance to nystatin, a polyene antibiotic (19). Although these mutants were viable there was speculation based on early growth experiments (20) that the *ERG6* step might be essential for the cell. If this were the case, it would make an excellent target for drug development since this step is not found in cholesterol biosynthesis thus eliminating one potential side effect problem. This possibility could not be fully explored until the *ERG6* gene was cloned and disrupted since the mutations originally isolated could be leaky and allow some ergosterol synthesis. The question of the essentiality of the *ERG6* gene was resolved by Gaber et al. (21) who showed

that *ERG6* disrupted strains were viable although loss of the *ERG6* gene product resulted in several altered phenotypes. Many of these phenotypes were consistent with altered physiological (22, 23) and membrane structural features (24, 25) that we have described previously in *erg6* mutants. Summarizing these findings, *erg6* mutants are compromised in several physiological areas (growth limitations on a variety of energy sources, mating difficulties, low transformation frequency, permeability alterations) due to the presence of membrane sterol that does not have the C-24 methyl group (21-25).

The most important of the alterations noted in *erg6* cells was the increased permeability of the cell to a variety of substances. This trait was demonstrated in our lab using a variety of molecules including dyes (22), cations (22) and spin labels used in electron paramagnetic resonance studies (24). Increased permeability was also employed in the cloning strategy (21) for the ERG6 gene and, subsequently, several other pathway genes. More recently, cloning of the LIS1 gene, mutations of which lead to increased uptake of lithium and sodium, was accomplished and sequencing revealed that this gene was identical to ERG6 (26). Our hypothesis is that disruption the ERG6 gene of C. albicans or inhibition of the ERG6 gene product in this organism would generate a phenotype similar to that seen in S. cerevisiae ERG6 disruptants. Such a phenotype would make the cell more sensitive to known antifungals and more permeable to inhibitory substances that would not normally be able to gain entry to the cell. We would also be interested in determining the effect of the erg6 mutation on the ability of C. albicans to form hyphae, a property necessary for pathogenesis. We have shown previously that a mutation in the ERG11 step seriously compromises the ability of this organism to form hyphae (27). With these possibilities in mind, we are seeking to disrupt both copies of the C. albicans ERG6 gene, test it for increased sensitivity to a variety of substances, and utilize the isolated ERG6 gene in a screen for the isolation of substances that block the function of the ERG6 gene product.

Most recently (28) we isolated and characterized the *S. cerevisiae* gene (*ERG25*) encoding the C-4 sterol methyl oxidase, the enzyme responsible for initiating the removal of the two methyl groups from the C-4 position. This work has leveraged our ability to isolate the *ERG25* gene from *C. albicans*. Since we have determined that the *ERG25* gene is essential to yeast for survival, verification of this phenotype in *C. albicans* might indicate that the *ERG25* gene product would be a good candidate target for the development of new antifungals. We now have the complete sequence of the *C. albicans ERG25* and are in the process of disrupting both copies to determine if it is essential in this organism as it is in *S. cerevisiae*.

We are also interested in the phenomenon, noted in *S. cerevisiae*, where essential genes in the sterol biosynthetic pathway can be suppressed by mutations in other genes and viability is thus restored. Defining such mechanisms will help determine and analyze such events should they occur in *C. albicans*.

<u>Background</u>: Sterols are required components of all eucaryotic membranes and numerous studies of the physiological and biophysical effects of alteration of membrane sterol content have indicated their importance for normal permeability functions(23-25). In addition, sterols play a critical role in maintaining the appropriate membrane viscosity such that membrane bound enzymes can perform normally (29). Our best information on ergosterol biosynthesis comes from studies using the common yeast, *S. cerevisiae*.

Sterols are synthesized as a separate branch of the isoprenoid pathway which also produces essential end products such as quinones, heme, and dolichols (30). In addition, intermediates such as farnesyl pyrophosphate and geranyl pyrophosphate are employed in the isoprenylation of proteins that require membrane anchoring for appropriate functioning (30).

The sterol branch of the isoprenoid pathway is shown in Figure 1. The first compound, squalene, is formed by the enzyme squalene synthase by combining two molecules of farnesyl pyrophosphate. The next two reactions yield lanosterol, the first sterol in the pathway. Mutations in any of the steps prior to lanosterol formation result in sterol auxotrophy and cells carrying such mutations must be provided exogenous sterol. Mutations in any of the three steps (*ERG11*, *ERG 24*, and *ERG25*) following lanosterol synthesis result in aerobic non-viability since cells producing sterol can no longer take up exogenous sterol and these mutants produce sterol intermediates that cannot be utilized in place of ergosterol(31, 32, 28). Mutations in the subsequent steps of the pathway have been shown to produce sterol intermediates capable of supporting growth. These mutants, however, do show altered phenotypes with those furthest from the ergosterol end product showing the most compromised characteristics.

Our knowledge of the sterol biosynthetic pathway in the pathogen, *Candida albicans*, is considerably less complete. The same basic steps are present although the order of some of the terminal reactions is altered compared to *S. cerevisiae*. The determination of essentiality for various steps in *S. cerevisiae* has been accomplished using gene disruption and allele replacement techniques. Since only a handful of genes in *C. albicans* have been cloned, much remains to learned about the essential reactions of the pathway in this organism.

In addition to characterization of the ergosterol biosynthetic steps in *C. albicans*, it is important that the phenomenon of genetic suppression be explored. Suppression occurs when a second mutation masks the effects of the primary mutation. In the case where the first mutation is lethal, the suppressor would result in viability. This is critical consideration if we are able to identify an essential reaction and design an inhibitor only to find that a suppressor mutation counteracts the inhibitor.

Suppression has been identified in all *ERG11*, *ERG24*, and *ERG25* steps of ergosterol biosynthesis in *S. cerevisiae*. Mutations in the *ERG11* gene are suppressed

by downstream mutations in the *ERG3* gene. This is possible because blocks at *ERG11* result in C-14 methylsterols that are acted upon by downstream enzymes. However, the enzyme encoded by the *ERG3* gene cannot complete the desaturation of the C-14 sterol intermediate resulting in the formation of a toxic sterol diol (33). Mutations in *ERG3* result in no diol formation and permit viability in *erg11* mutants (34). Mutations in the *ERG24* gene result in the accumulation of ignosterol, a sterol that cannot support growth. However, a mutation, *fen1* (35), can suppress *erg24* and permit the cell to grow on ignosterol. The nature of *fen1* remains undefined. Finally, we have recently reported (36) on a novel mechanism of suppression of a mutant of the *ERG25* gene. This suppression involves two separate mutations. One is a mutation in *ERG11* while the second is a leaky mutation in one of the heme biosynthetic genes resulting in low levels of heme.

Suppression in the sterol pathway has not been explored in *C. albicans*. It is curious to note that *ERG11* mutations in *C. albicans* are viable and suppression is not required to permit growth. This means that *C. albicans* can either tolerate the sterol diol that accumulates (31) or that this organism is able to selectively sequester this molecule. Thus, sterol requirements and suppression may vary between *S. cerevisiae* and *C. albicans*. Both of these parameters will be addressed during the course of our studies.

BODY

Experimental Methods and Results:

CLONING AND SEQUENCING OF THE CANDIDA ALBICANS ERG6 GENE

Previously, we reported that the *C. albicans ERG6* gene was cloned using a *Candida* gene library transformed into a *S. cerevisiae erg6* strain. We obtained two complementing and overlapping clones with inserts of 7.8 kb and 14 kb. pIU885 contains a 2.4 kb *XbaI-EcoRI Candida* DNA fragment which is able to complement the *Saccharomyces erg6* mutation (Figure 2). This DNA fragment was sequenced using the Sanger dideoxy chain termination method and the DNA and amino acid sequences are presented in Figure 3. The *Candida ERG6* gene encodes the sterol methyl transferase which contains 377 amino acids and is 66% identical to the *Saccharomyces* enzyme. Figure 4 shows an alignment between the *Candida*, *Saccharomyces*, *Arabidopsis*, and *Triticum* sterol methyl transferases and the percent identity of *Candida* to the latter two are 40% and 49%, respectively. A 9 amino acid region (Figure 4; amino acids 129-137) represents the highly conserved S-adenosyl methionine binding site (37).

DISRUPTION OF THE ERG6 GENE

Disruption of the *Candida ERG6* gene to derive a sterol methyltransferase deficient strain was made more difficult since *Candida*, unlike *Saccharomyces*, is an obligate diploid and thus both copies of the *ERG6* gene must be disrupted. To

accomplish this, we used the "ura blaster" system developed by Fonzi (38). Essentially the ura blaster contains ~1.2 kb repeat elements of *hisG* (derived from Salmonella) flanking the Candida URA3 gene (see Figure 2). A plasmid containing the ura blaster inserted into the ERG6 gene is shown in Figure 2. The 2.4 kb XbaI-Ecol ERG6 DNA fragment was cloned into the Bluescript vector pKS(+) in which a *HindIII* site was filled in with the Klenow fragment of DNA polymerase I (pIU886). pIU886-L was subsequently derived by deleting a 0.7 kb HindIII within the ERG6 coding sequence, filling in this site with Klenow followed by the addition of BamHI linkers. Plasmid 5921, containing the ura blaster, was digested with SnaBI and StuI, both blunt cutting enzymes, followed by religation. This resulted in a deletion of 6 bp in one of the hisG regions and destruction of these two sites. The modified 5921 plasmid was then digested with BamHI and BglII to release the 3.8 kb ura blaster which was then ligated into pIU886-L that had been digested with BamHI to generate pIU887-A, a plasmid containing the ura blaster inserted into ERG6 (Figure 2).

Candida strain CAI4 was transformed using the 5.3 kb *BgIII-Sna*BI fragment containing the ura blaster and *ERG6* flanking recombinogenic ends of 0.8 and 0.9 kb. Transformants containing the single disrupted *ERG6* allele resulting in heterozygosity for *ERG6* was confirmed using PCR after selection for loss of the *URA3-hisG* region. Intrachromosomal recombination between the linear *hisG* sequences results in loss of one of these *hisG* repeats and the *URA3* thus permitting reuse of the ura blaster for the subsequent disruption of the *ERG6* gene on the homologous chromosome. Selection for colonies on 5-Fluoro-orotic acid results in growth of only uracil requiring strains (39).

CREATION OF A HOMOZYGOUS erg6 STRAIN

The creation of a *Candida erg6* mutant strain in which both alleles were disrupted was accomplished in two different ways. The *ERG6* heterozygote was placed onto plates containing high concentrations of nystatin ($20\mu g/ml$) and after 3 days nystatin resistant colonies appeared. We surmised that mitotic recombination might result in homozygous *ERG6* and *erg6* segregants and these nystatin resistant colonies might be the *erg6* homozygotes. When colony purified, these resistant colonies indeed turned out to be *erg6* homozygotes (see next section). The second method used to generate *erg6* homozygotes was to transform the *ERG6* heterozygote with the ura blaster. This time two kinds of transformants were obtained -wild type and slow growing colonies. Both types of colonies were tested for resistance to nystatin and only the slower growing colonies appeared to be nystatin resistant.

CONFIRMATION OF erg6 HOMOZYGOSITY

All putative *erg6* mutants generated by both methods were nystatin resistant. The sterols isolated from wild type and putative *erg6* homozygotes were analyzed by UV spectrophotometry and gas-chromatography/mass spectroscopy. Yeast sterol

samples were scanned by UV in the 200-300nm range-for absorption maxima at 262, 271, 282, and 293 indicating a conjugated diene system in the B ring of the sterol molecule. Additionally, *erg6* mutants accumulate sterols with a conjugated diene system in the sterol side-chain resulting in absorption maxima at 230 and 238nm (19). All of our putative *erg6* homozygotes contained *erg6*-like sterol profiles. Additionally, GC/MS of *erg6* mutant sterols confirmed that only cholesterol-like (C-27) sterols accumulate since the side-chain cannot be methylated. Figure 5 represents a GC profile demonstrating that the putative *erg6* mutants accumulate C-27 sterols and are deficient in side-chain transmethylation. Whereas the predominant sterol in the CAI4 wild type is ergosterol (peak B, 76%,), the principal sterols in *erg6* mutants are: zymosterol (peak A, 43%), cholesta-5,7,24-trien-3β-ol (peak D, 6%), cholesta-7,24-dien-3β-ol (peak E, 9%), and cholesta-5,7,22,24-tetraen-3β-ol (peak F, 29%). Lastly, PCR analyses using combinations of four different primers within the *ERG6* region and the *hisG* region confirmed that both *ERG6* wild type alleles were disrupted.

CLONING AND SEQUENCING OF THE CANDIDA ALBICANS ERG25 GENE

Previously, we reported that the *C. albicans ERG25* gene was cloned using a *Candida* gene library transformed into a *S. cerevisiae erg25* strain. Again, we obtained two complementing and overlapping clones with inserts of 6 kb and 3.8 kb. Plasmid pIU870 (see Figure 6) containing a 2.9 kb *BamHI-BgIII* DNA fragment inserted into pRS316 complemented the *erg25* mutation. Two subclones of pIU870 containing a 0.8 kb *Hind* III fragment (pIU875) and a 0.6 kb *Hind* III-*BgIII* fragment (pIU876) were used for DNA sequencing (Figure 6). A third subclone containing a 1.5 kb *BgIII-Eco*RI fragment was found not to contain *ERG25* sequence but instead DNA encoding isocitrate dehydrogenase. Figure 7 shows the DNA and amino acid sequences of *Candida ERG25*. The *Candida* sterol methyl oxidase is a 294 amino acid protein having 65% identity with the *Saccharomyces* enzyme and 36% identity with the human enzyme (see Figure 8). Conserved sequences among all three enzymes are three histidine clusters required for binding of non-heme iron (Figure 7, boxed region). Additionally, all three enzymes contain the ER retrieval motif KKXX or KXKXX required for retrieving protein from the Golgi back to the ER (40).

Discussion and Conclusion:

The project has three aims as shown in the Statement of Work (see Appendix). The first aim seeks to explore the *ERG6* gene of *C. albicans* as a potential target for the development of new antifungals. One year ago we reported the cloning of this gene by complementation of an *erg6* mutant of *S. cerevisiae* with a *Candida* library. Sequencing of the *Candida ERG6* gene is now complete (Figure 3). We have also completed disruption of both alleles in this organism and found the *ERG6* to be non-essential for viability. We are now in the process of defining the characteristics of cells harboring the double *ERG6* disruption and preliminary data indicate that removal of *ERG6* gene function results in cells with altered growth

indicate that removal of *ERG6* gene function results in cells with altered growth characteristics and permeability characteristics similar to those described for *erg6* mutants of *S. cerevisiae*.

The second aim of our project seeks to isolate and disrupt the genes responsible for the C-4 demethylation step in ergosterol biosynthesis. This is the final unexplored step in fungal sterol pathway. This effort is tied to our work in *S. cerevisiae* where the initial experimentation must be done. The first phase in C-4 demethylation is an oxidation of the C-4 methyl group. The gene (*ERG25*) responsible for this reaction (sterol methyl oxidase) has been isolated and characterized in our laboratory (28). The *Candida ERG25* has been isolated by complementation of a *S. cerevisiae erg25* mutant with a *Candida* library. Since our last report the *Candida ERG25* gene has been completely sequenced (Figure 7). We are now in the process of disrupting both alleles of *ERG25* in this organism. This is a more difficult process since it is likely that the *erg25* phenotype will not be viable and *Candida* does not take up exogenous sterol to maintain viability. We are devising some protocols that will allow us to perform the disruptions in a background that will permit viability.

The final aim of our project is to explore the phenomenon of suppression in *Candida*. As is the case in the C-4 demethylation aim, we will first define suppression mechanisms in *S. cerevisiae* since this organisms has many genetic and physiological advantages. We have recently described a unique suppression mechanism for *erg25* mutants of *S. cerevisiae* and will investigate suppression of this step in *Candida* once we have created the double disruption.

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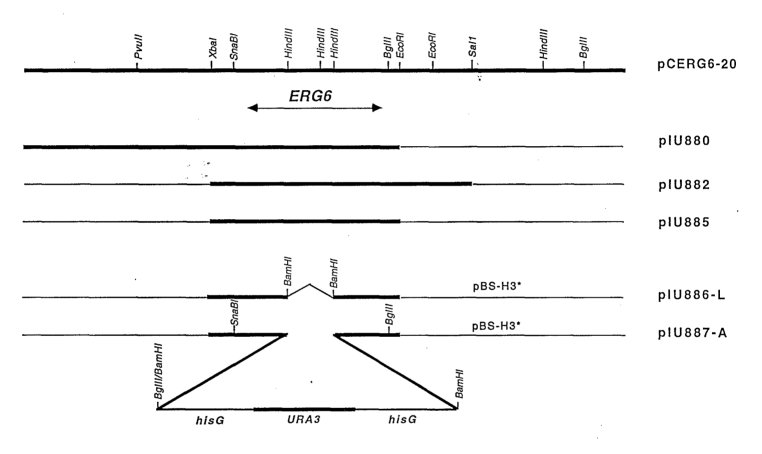
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Figure 1. The ergosterol biosynthetic pathway from lanosterol to ergosterol.





1 kb

Figure 2. A *Candida albicans ERG6* genomic clone (pCERG6-20) with restriction sites and three complementing subclones, pIU880, pIU882, and pIU885. Modification of pIU885 to pIU886-L and subsequent disruption with the ura blaster shown in pIU887-A are represented.

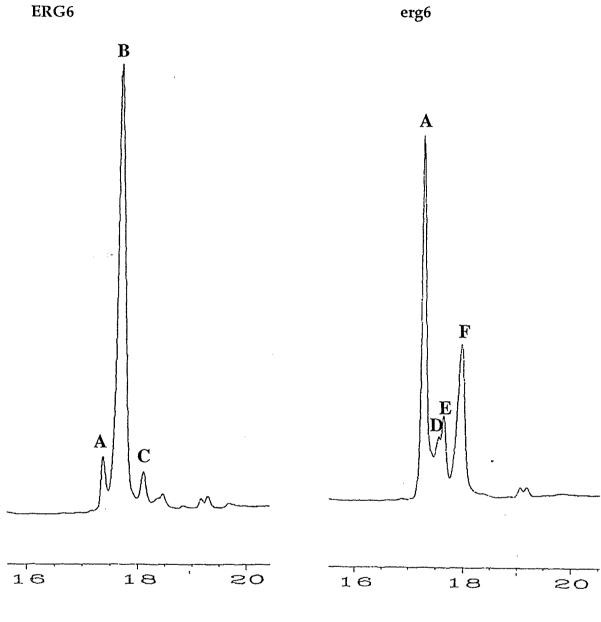
1 80	80 atttaatttagaaattttcaacctaattaattcataatctaagattcaactcattaaca ATG TCT CCA GTT CAA																				
																М	S	P	V	Q	5
155		GCA A	GAA E	AAA K	AAT N	TAC Y	gaa E	AGA R	GAT D	GAA E	CAA Q	TTC F	ACT T	AAA K	GCT A	TTA L	CAT H	GGT G	GAA E	TCT S	25
216	TAT Y	AAA K	AAA K	ACT T	GGG G	TTA L	TCA S	GCT A	TTA L	ATA I	GCT A	AAA K	TCT S	AAA K	GAT D	GCT A	GCT A	TCT S	GTT V	GCT A	45
277	GCT A	GAG E	GGT G	TAT Y	TTC F	AAA K	CAT H	TGG W	GAT D	GGT G	GGT G	ATT I	TCT S	AAA K	GAT D	GAT D	GAA E	GAG E	AAA K	AGA R	65
338	TTG L	AAT N	GAT D	TAT Y	TCC S	CAA Q	TTG L	ACT T	CAT H	CAT H	TAT Y	TAT Y	AAT N	TTA L	GTC V	ACT T	GAC D	TTT F	TAT Y	GAA E	85
399	TAT Y	GGT G	TGG W	GGT G	TCT S	TCA S	TTC F	CAT H	TTT F	TCA S	AGA R	TAT Y	TAT Y	AAA K	GGT G	GAA E	GCT A	TTT F	AGA R	CAA Q	105
460	GCT	ACT	GCT	AGA	CAT	GAA	CAT	TTC	TTG	GCC	CAT	AAA	ATG	AAT	CTT	ААТ	GAA	AAC	ATG	AAA	
	A	Т	A	R	Н	E	Н	F	L	A	Н	K	М	N	L	N	Е	N	М	K	125
521	GTT V	TTA L	GAT D	GTT V	GGT G	TGT C	GGT G	GTA V	GGT G	GGT G	CCT P	GGT G	AGA R	GAA E	ATC I	ACA T	AGA R	TTT F	ACT T	GAT D	145
582	TGT C	~GAA E	ATT I	GTT V	GGA G	TTA L	AAT N	AAT N	AAT N	GAT D	TAT Y	CAA Q	ATT I	GAA E	AGA R	GCT A	aat N	CAT H	TAT Y	GCT A	165
643	AAA K	AAA K	TAC Y	CAT H	TTA L	GAT D	CAT H	AAA K	TTA L	TCT S	TAT Y	GTT V	AAA K	GGT G	GAT D	TTT F	ATG M	CAA Q	atg M	GAT D	185
704	TTT F	GAA E	CCA P	GAA E	TCA S	TTC F	GAT D	GCT A	GTT V	TAT Y	GCC A	ATT I	GAA E	GCT	ACC T	GTT V	CAT H	GCT A	CCA P	GTT V	205
765	TTA L	GAA E	GGA G	GTT V	TAT Y	TCA S	GAA E	ATT I	TAT Y	AAA K	GTT V	TTG L	AAA K	CCA	GGT G	GGT G	ATT I	TTC F	GGT G	GTT V	225
826	TAT Y	GAA E	TGG W	GTC V	ATG	ACT T	GAT D	AAA K	TAC Y	GAT D	GAA E	ACT T	AAT N	GAA E	GAA E	CAT H	CGT R	'AAA K	ATT I	GCT A	245
887	TAT Y	GGT G	' ATT	GAA	GTC V	GGT G	GAT	GGT G	ATT I	CCA	AAA K	ATG	TAT Y	TCT	CGT R	AAA K	. GTI V	GCT A	GAA E	CAA Q	265
948	GCT A	TTG	AAA K	raa <i>i</i> N	GTT	GGA	. TTI F	GAA	ATT	GAA	TAT Y	CAA Q	. AAA	A GAT	TTG	GCT	GAT	GTT	GAT		285
1009					TAT Y													r TTT F	GGT G	GAT D	305
1070			_					٠								_			_	GGT	303
	Y	L	T	V	F	R	T	s	R	I	G	R	F	I	T	T	E	s	V	G	325
1131	TTA L	ATO M	G GAZ	A AAZ	A ATT	r GGT G	TTA L	A GCT	P CCA	A AAA K	G G	TCT S	r aaz K	A CAZ Q	V GTT	T ACT	CAT H	r gct A	TTA L	GAA E	345
1192	GAT D	r gc	r gc	r gt	r aa: N	r TT	A GT	F GAZ	A GGT G	r GG?	r aga	Q CAZ	A AA	A TTO	F TT	r AC	r cci	A ATO	G ATO	TTG L	365
1253	TAC Y	C GT	r gr	r ag. R	A AA K	A CC	A TT	A GAZ	A AAG	G AAZ K	A GAT	Г ТА <i>і</i> *	A tg	gggc	ttga	caaa	çaac	aagt	aaggt	gagtt	377
1322 1402	22 tatgttgggggtgttcaattcgtgtatctattcatagaggtatttgatttgcaatttgttttttgtttattctatttatt									-											

Figure 3. The DNA and amino acid sequences of the Candida albicans ERG6 gene.

```
MSPVQLAEK-NYERDEQFTKALHGESY-KKTGLSAL
C. a. ERG 6
          1
S. c. ERG 6
             MSETELR - - - - KRQAQFTRELHGDDIGKKTGLSAL
             MDSLTLFFTGALVAVGIY-WFLCVLGPAERKGKRAV 35
MFVFCLCTRCRICRVSSFPVLLLFMFIHLSYFFLVL 36
A. t. ERG 6
T. a. ERG 6
             IAKSKDAASVAAEGYFKHWDGGISKDDEEKRLNDYS 70
C. a. ERG 6
         35
S. c. ERG 6
             M S K N N S A Q K E A V Q K Y L R N W D G R T D K D A E E R R L E D Y N
         32
             DLSGGSISAEKVQD<u>N</u>YKQYW<u>S</u>FFRRPK<u>EIET</u>AEK<u>V</u>P
A. t. ERG 6
             LLILGQFFFTRYEKYHGYYGGK----EESRKSNYT
         37
T. a. ERG 6
             QLTHHYYNLVTDFYEYGWGSSFHFSRYYKGEAFRQA 106
C. a. ERG 6
         71
             EATHSYYNVVTDFYEYGWGSSFHFSRFYKGESFAAS
         68
                                                                    103
S. c. ERG 6
             DFVDTFYNLVTDIYEWGWGQSFHFSPSIPGKSHKDA 107
DMVNKYYDLATSFYEYGWGESFHFAHRWNGESLRES 103
         72
A. t. ERG 6
T. a. ERG 6
        107
             TARHEHFLAHKMNLNE<u>NMK</u>VLDVGCGVGGPGREITR 142
C. a. ERG 6
             IARHEHYLAYKAGIQRGDLVLDVGCGVGGPAREIAR
S. c. ERG 6
        104
             TRLHEEMAVDLIQVKPGQKILDVGCGVGGPMRAIAS
        108
A. t. ERG 6
                                                                    143
             IKRHEHFUALQLELKPGMKVLDVGCGIGGPLREIAR 139
        104
T. a. ERG 6
        143
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C. a. ERG 6
             FTGCNVIGLNNNDYOIAKAKYYAKKYNLSDQMDFVK 175
HSRANVVGITINEYOVNRARLHNKKAGLDALCEVVC 179
FSSTSVTGLNNNDYOITRGKALNRSVGLGATCDFVK 175
        140
S. c. ERG 6
A. t. ERG 6
        144
T. a. ERG 6
        140
             GDFMQMDFEPESFDAVYALEATVHAPVLEGVYSEIY
C. a. ERG 6
        179
             GDFMKMDFEENTFDKVYAIEATCHAPKLEGVYSEIY
GNFLQMPFDDNSFDGAYSIEATCHAPKLEEVYAEIY
S. c. ERG 6
        176
                                                                    211
        180
A. t. ERG 6
                                                                    215
        176
              ADFMKMPESDNTFDAVYAIEATCHAPDPVGCYKEIY 211
T. a. ERG 6
             KVLKPGGIEGVYEWVMTDKYDETNEEHRKTAYGIEV 250
        215
C. a. ERG 6
             KVLKPGGTFAVYEWVMTDKYDENNPEHRKIAYEIEL 247
RVLKPGSMYVSYEWVTTEKFKAEDDEHVEVIQGIER 251
        212
S. c. ERG 6
A. t. ERG 6
        216
              RVLKPGQCFAVYEWCITDHYDPNNATHKRIKDEIEL 247
T. a. ERG 6
        212
             GDGIPKMYSRKVAEQALKNVGFEIEYQKDLADVDDE
GDGIPKMFHVDVARKALKNCGFEVLVSEDLADNDDE
        251
C. a. ERG 6
                                                                    286
        248
S. c. ERG 6
             GDALPGLRAYVDIAETAKKVGFEIVKEKDLASPPAE 287
GNGLPDIRSTRQCLQAVKDAGFEVIWDKDLAE-DSP 282
        252
A. t. ERG 6
T. a. ERG 6
        248
              IPWYYPLSGDLKFCQTFGDYLTVFRTSRIGRFITTE 322
        287
C. a. ERG 6
              IPWYYPLTGEWKYVQNLANLATFFRTSYLGRQFTTA 319
S. c. ERG 6
         284
A. t. ERG 6
        288
              LPWYLPL - DPSRFS - - - - - LSSERLTTVGRIITRN 311
T. a. ERG 6
        283
              SVGLMEKIGLAPKGSKQVTHALEDAAVNLVEGGRQK 358
         323
C. a. ERG 6
              MVTVMEKLGLAPEGSKEVTAALENAAVGLVAGGKSK 355
S. c. ERG 6
         320
                 QILSAVGVAPKGTVDVHEMLFKTADCLTRGGETG 341
        306
A. t. ERG 6
              MVKVLEYVGLAPEGSQRVSSFLEKAAEGLVEGGKKE 347
T. a. ERG 6
        312
              LFTPMMLYVVRKPLEK
         359
C. a. ERG 6
                                                                     374
              LFTPMMLFVARKPENAETPSQTSQEATQ
         356
S. c. ERG 6
                                                                     383
         342
              IESPMHMILCRKPESPEESS
A. t. ERG 6
                                                                     361
              IFTPVYFFVVRKPLSE
T. a. ERG 6
         348
                                                                     363
```

Figure 4. Alignment of the amino acid sequences of the sterol methyl transferases from *Candida albicans* (C. a.), *Saccharomyces cerevisiae* (S. c.), *Arabidopsis thaliana* (a. t.), and *Triticum ativum* (t. a.).

UNPUBLISHED DATA



RETENTION TIME

Figure 5. Gas chromatography wild type and an *erg6* strain of *Candida albicans*. Peak A; zymosterol, Peak B; ergosterol, Peak C; fecosterol, Peak D; cholesta-5,7,24-trien-3β-ol, Peak E; cholesta-7,24-dien-3β-ol, Peak F; cholesta-5,7,22,24-tetraen-3β-ol.

ERG25 Clones

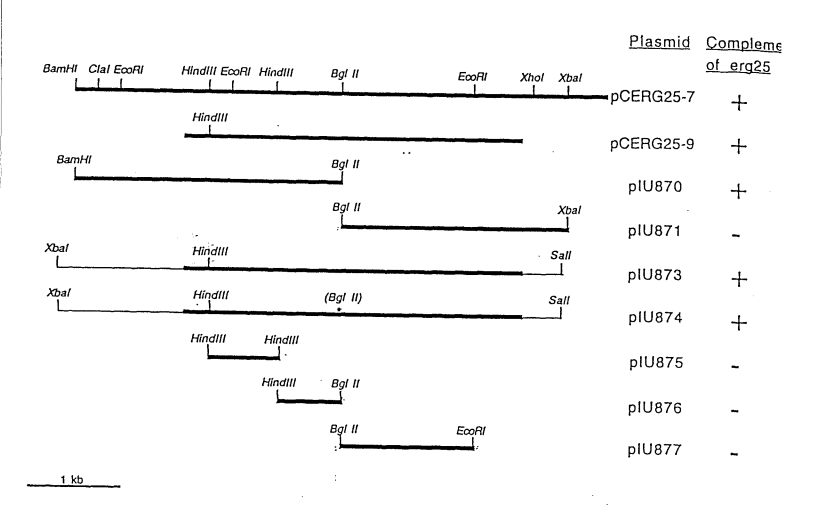


Figure 6. Restriction endonuclease sites on the *Candida albicans ERG25* genomic clones, pCERG25-7 and pCERG25-9. The ability of subclones pIU870 through pIU877 to complement an *erg25* recipient strain is indicated in the right column. Bold lines represent genomic DNA and thin lines represent plasmid pRS316 DNA.

1 80	tttt tatt	gatt tatt	catt tgtt	aatt tgtt	gtta tgaa	tatt gttt	tcaa ata	ATG	TCT	TCC	ATT	AGT	AAT	GTT	TAT	CAT	GAC	TAT	TCG	AGT	
								M	S	S	I	S	N	V	Y	Н	D	Y	S	S	13
147	TTT F	CTG S	AAT N	GCA A	ACT T	ACT T	TTT F	TCC S	CAA Q	GTT V	TAT Y	CAA Q	AAT N	TTC F	TAA N	CAA Q	TTA L	GAT D	AAT N	TTA L	33
208	AAT N	gtt V	TTT F	GAA E	AAA K	TTA L	TGG W	GGG G	TCA S	TAT Y	TAT Y	TAT Y	TAT Y	ATG M	GCC A	AAT N	GAT D	TTA L	TTT F	GCT A	53
269	ACT T	GGA G	TTA L	TTA L	TTT F	TTT F	TTA L	ACT T	CAT H	GAA E	TTA I	TTT F	TAT Y	TTT F	GGT G	AGA R	TGT C	TTA L	CCA P	TGG W	73
330	GCT A	ATA I	ATT I	GAT D	AGA R	ATT I	CCT P	ТАТ Ү	TTT F	AGA R	AAA K	TGG W	AAA K	ATT I	CAA Q	GAT D	GAA E	AAA K	ATC I	CCT P	93
391	AGT S	GAT D	AAA K	GAA E	CAA Q	TGG W	gaa E	TGT C	CTT L	aaa K	TCA S	GTT V	TTA L	ACA T	TCT S	CAT H	TTC F	TTA L	GTT V	GAA E	113
452	GCT A	TTC F	CCA P	ATT I	TGG W	TTT F	TTC F	CAT H	CCA P	TTA L	TGT C	CAA Q	AAA K	ATT I	GGT G	ATT I	AGT S	TAT Y	CAA Q	GTA V	133
513	CCA P	TTC F	CCT P	AAA K	ATT I	ACT T	GAT D	ATG M	TTG L	ATT I	CAA Q	TGG W	GCA A	GTA V	TTT F	TTT F	GTT V	TTG L	GAA E	GAT D	153
574	ACT T	TGG W	CAT H	TAT Y	TGG W	TTT F	CAT H	AGA R	GGA G	TTA L	CAT H	TAT Y	GGG G	GTT V	TTC F	TAT Y	AAA K	TAT Y	ATT I	CAT H	173
635	AAA K	CAA Q	CAT H	CAT H	AGA R	TAT Y	GCT A	GCT A	CCA P	TTT F	GGA G	TTG L	GCA A	GCA A	GAA E	TAT Y	GCT A	CAT H	CCA P	GTT V	193
696	GAA E	GTT V	GCC A	TTA L	TTA L	GGA G	TTG L	GGT G	ACG T	GTT V	GGT G	ATT I	CCG P	ATT I	GTT V	TGG W	TGT C	CTT L	ATC I	ACT T	213
757	GGT G	AAC N	TTG L	CAT H	CTT L	TTC F	ACA T	GTT V	TCC S	ATT I	TGG W	ATC	ATT	TTA	AGA R	TTA	TTC F	CAA Q	GCC A	GTT V	233
818	GAT D	GCT A	CAT	TCC S	GGT G	TAT Y	GAA E	TTC F	CCT P	TGG W	TCT S	TTA L	CAT H	raa n	TTC F	TTG	CCA P	TTT F	TGG W	GCT A	253
879	GGT G	GCT A	GAT	' CAT	CAT	' GAT D	GAA	CAT H	CAT H	H CAT	TAT Y	TTC F	: ATI		' GGA	TAC Ý	TCT S	TCA S	TCI S	TTT F	273
940	AGA R		TGG W	GAT D	TTC F	ATT	TTG	GAT D	' ACC	GAA	GCT A	G G	CCA P	A AAA K	A GCT		AAG K	GGT G	'AGA	GAA E	293
1001	GAC D	AAA K	GTC V	: AAA	CAA Q	raa <i>l</i> N	GTI V	GAA	AAA K	TTA L	CAA Q	AAC K	AAC K	AAC N	TT#	A TAG	aga	ıgaga	ıaaga	ıgtat	308
1065 1145																					

Figure 7. The DNA and amino acid sequences of the Candida albicans ERG25 gene.

C. a. ERG 25	1	M S S I S N V Y H D Y S S F S N A T T F S Q V Y Q N F N Q L D M S A V F N N A T L S G L V Q A S T Y S Q T L Q N V A H Y Q P M A T N E S V S I F S S A S L A V E Y V D S L L P	31
S. c. ERG 25	1		31
H. s. ERG 25	1		25
C. a. ERG 25	32	N L N V F E K L W G S Y Y Y Y M A N D L F A T G L L F F L T H E Q L N F M E K Y W A A W Y S Y M N N D V L A T G L M F F L L H E E N P L Q E P F K N A W N Y M L N N Y T K F Q I A T W G S L I V H E	63
S. c. ERG 25	32		63
H. s. ERG 25	26		59
C. a. ERG 25	64	I F Y F G R C L P W A I I D R I P Y F R K W K I Q D E K I P S D K E F M Y F F R C L P W F I I D Q I P Y F R R W K L Q P T K I P S A K E A L Y F L F C L P G F L F Q F I P Y M K K Y K I Q K D K P E T W E N	97
S. c. ERG 25	64		97
H. s. ERG 25	60		93
C. a. ERG 25	98	QWECLKSVLTSHFLVEAFPIWFFHPLCQKIGUSY	131
S. c. ERG 25	98	QLYCLKSVLLSHFLVEAIPIWTFHPMCEKLGITV	131
H. s. ERG 25	94	QWKCFKVLLFNHFCIQLPLICGTYYFTEYFNIPY	127
C. a. ERG 25	132	QVPFPKITDMLIQWAVFFVLEDTWHYWFHRGEVPFPSLKTMALEIGLFFVLEDTWHYWAHRLDWERMPRWYFLLARCFGCAVIEDTWHYFLHRL	162
S. c. ERG 25	132		162
H. s. ERG 25	128		159
C. a. ERG 25	163	LHYGVFYKYIHKQHHRYAAPFGLAAEYAHPVEVA	196
S. c. ERG 25	163	FHYGVFYKYIHKQHHRYAAPFGLSAEYAHPAETL	196
H. s. ERG 25	160	LHHKRIYKYIHKVHHEFQAPFGMEAEYAHPLETL	193
C. a. ERG 25	197	L L G L G T V G I P I V W C L I T G N L H L F T V S I W I I L R L F S L G F G T V G M P I L Y V M Y T G K L H L F T L C V W I T L R L F I L G T G F F I G I V L L C D H V I L L W A W V T I R L L	230
S. c. ERG 25	197		230
H. s. ERG 25	1 94		222
C. a. ERG 25	231	QAVDAHSGYEFPWSLHNFLPFWAGADHHDEHHHY	264
S. c. ERG 25	231	QAVDSHSGYDFPWSLNKIMPFWAGAEHHDLHHHY	264
H. s. ERG 25	223	ETIDVHSGYDIPLNPLNLIPFYAGSRHHDFHHMN	256
C. a. ERG 25	265	FIGGYSSSFRWWDFILDTEAGPKAKKG FIGNYASSFRWWDYCLDTESGPEAKASREERMKK FIGNYASTFTWWDRIFGTDSQYNAYNEKRKKFEK	291
S. c. ERG 25	265		298
H. s. ERG 25	257		290
S. c. ERG 25	299	R A E N N A Q K K T N	309
H. s. ERG 25	291	K T E	293

Figure 8. Alignment of the amino acid sequences of the C-4 sterol methyl oxidases from Candida albicans (c. a.), Saccharomyces cerevisiae (S. c.), and human (H. s.).

<u>Appendix</u>

STATEMENT OF WORK

- Aim 2 Cloning and disruption of the C-24 Transmethylase gene (ERG6) of Candida albicans
 - -cloning by complementation of a *C. albicans* genomic library with a *Saccharomyces cerevisiae erg6* mutant
 - -confirmation of plasmid-borne phenotype (FOA,) GC/MS analysis
 - -characterization by restriction mapping and subcloning
 - -determination of essentiality by sequential disruption
 - -physiological characterization of *C. albicans ERG6* disruptions including susceptibility testing

Months 0-24

Aim 1 Isolation of C-4 demethylase mutants of *C. albicans*Following isolation of the three genes for C-4 demethylation from *S. cerevisiae*:

Isolation of C-4 demethylase genes from C. albicans

- -complementation of *S. cerevisiae* C-4 demethylase mutants with a genomic library from *C. albicans*
- -confirmation of plasmid-borne phenotype (FOA), GC/MS analysis
- -characterization by restriction mapping and subcloning
- -gene disruption and allele replacement (sequential)
- -analysis of essentiality
- sequencing of the C. albicans C-4 demethylase genes

Months 12-36

Aim 3 Suppressor analysis

isolation of suppressors of C4 demethylase mutants characterization of suppressors
GC/MS analysis
sensitivity to inhibitors
Months 30-48



DEPARTMENT OF THE ARMY

US ARMY MEDICAL RESEARCH AND MATERIEL COMMAND 504 SCOTT STREET FORT DETRICK, MARYLAND 21702-5012

7/19/2000

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6 Jul 00

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